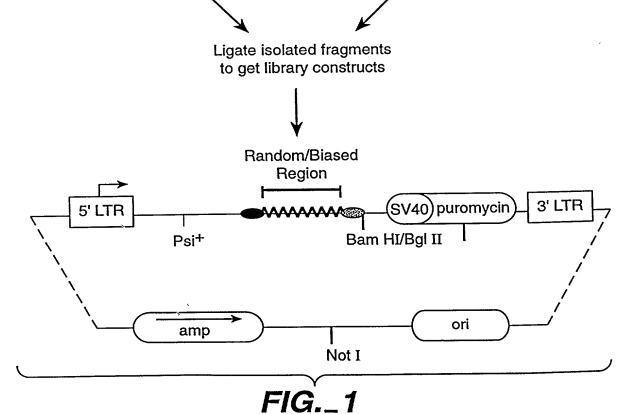
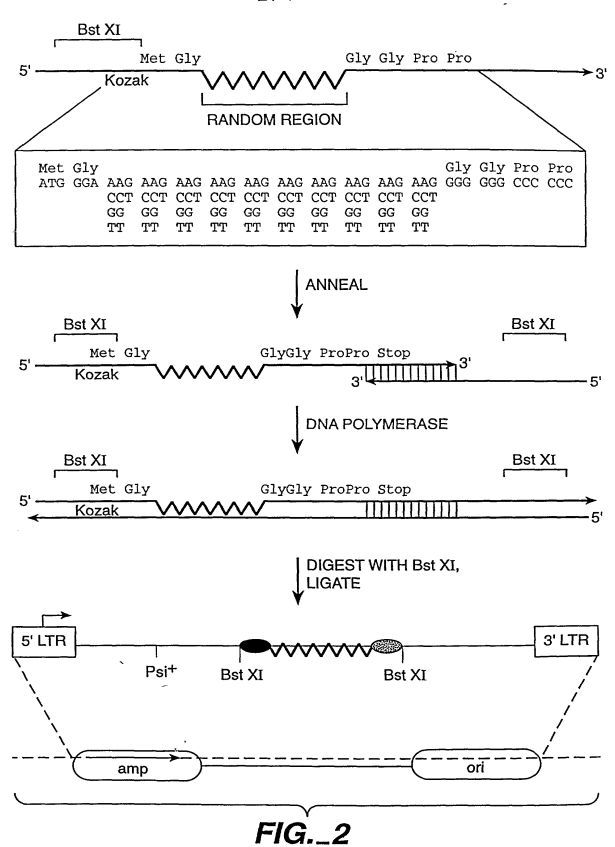
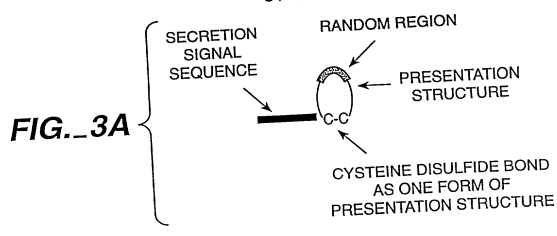
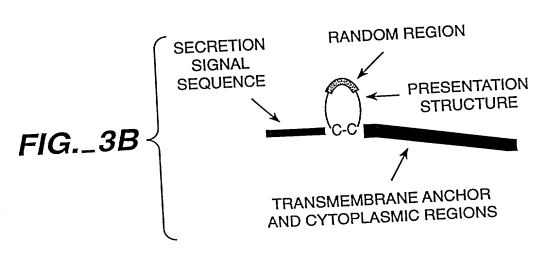


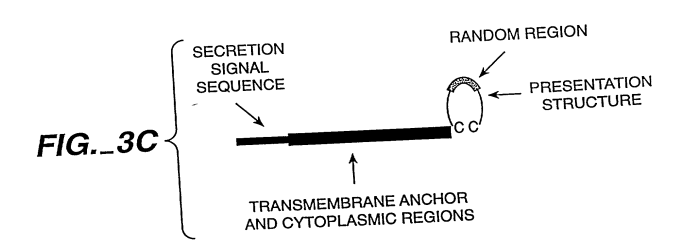
- PCR pBabe-puro with Primers 1 and 2.
- Cut PCR with Bgl $\scriptstyle\rm II$ and Not $\scriptstyle\rm I.$
- Isolate PCR fragment(s) containing randomized region from primer 1.
- Cut pBabe-puro with Bam HI and Not I.
- Isolate Bam/Not fragment with SV40-puro, 3'LTR, and bacterial ori.











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